I. Subjective Data

- A. Refrain from providing in the following conditions
 - 1. Undiagnosed, abnormal vaginal bleeding
 - 2. Progestagen-dependent tumor
 - 3. Active venous thromboembolic disorder
 - 4. Presence or history of severe hepatic disease, as long as liver function values have not returned to normal; active viral hepatitis.
 - 5. Hypersensitivity to the active substance or to any of the excipients of Implanon.
 - 6. Breast cancer current or within the last 5 years
 - 7. Sustained hypertension that develops during the use of Implanon discontinue use.
- B. Exercise caution in the following situations and carefully monitor for adverse effects. Women in this category who choose to use Implanon, where the clinician/physician determines that Implanon can be used, must be provided with information regarding the way in which these conditions may add a health risk for her. This discussion must be documented.
 - 1. Pre-existing breast cancer and no recurrence in the last 5 years.
 - 2. History of thromboembolic disorders
 - 3. Use of certain medications may make Implanon less effective, specifically those that induce cytochrome P450 enzymes resulting in an increased clearance of sex hormones in the liver. Some examples are: phenytoin, carbamazepine, phenylbutazone, barbiturates, the herbal remedy St. John's Wort, rifampin/rifampicin, griseofulvin (see Precautions). Women in long-term treatment with these drugs should consider another method of birth control. Please see package insert for complete list. Recommend concomitant use of a barrier method if use of these medications is short term.
 - 4. Cardiovascular conditions such as sustained hypertension with or without vascular disease, current or history of ischemic heart disease, history of CVA.
- C. Advantages outweigh theoretical or proven disadvantages
 - 1. Severe headaches including migraine with or without focal neurologic symptoms.
 - 2. Diabetes with or without vascular disease, nephropathy, retinopathy, neuropathy carefully monitor during first few months.

II. Objective Data

- A. Physical exam as per Title X regulations (follow package insert).
- B. Laboratory testing as per Title X regulations.

III. Assessment/Plan

- A. Client Education/Informed Consent
 - 1. Client must sign the Implanon insertion consent and information sheet.
 - 2. Client will be informed that Implanon offers no protection against sexually transmitted diseases; she should be advised to use condoms if she has concerns about potential exposure. There will be documentation of this in the client record.
 - 3. Certain medications may make Implanon less effective, specifically those that induce cytochrome P450 enzymes resulting in an increased clearance of sex hormones in the liver. Some examples are: phenytoin, carbamazepine, phenylbutazone, barbiturates, the herbal remedy St. John's Wort, rifampin/rifampicin, griseofulvin (see package insert for complete list). Women in long-term treatment with these drugs should consider another method of birth control.
 - 4. Client will be informed that if she wishes to discontinue the Implanon, she should make an appointment at the clinic for removal. If she does not wish to become pregnant she must start using another method on the day of removal.
- B. Insertion of Implants

Implanon may be inserted:

- 1. within the first 1 5 days of a regular menses;
- 2. at any time, if the client is currently using a highly effective method of birth control;
- 3. immediately post-abortion;
- 4. within 21 28 days after delivery or after a second trimester abortion
 - a. A prudent approach is to wait until breastfeeding is established at 4-6 weeks postpartum.
- beyond 28 days or if intercourse has already occurred, once pregnancy is ruled out.

IV. Follow-Up

- A. The client may return for an insertion site check if she has concerns about the Implanon site.
- B. Client should return to clinic at three months to have an evaluation of her satisfaction with the method. At this visit client should fill in the Hormonal Evaluation form. She may have her blood pressure checked, and be weighed if she wishes.
- C. Clients should be advised to call the clinic for an appointment for any of the following:
 - 1. arm pain; pus or bleeding at the insertion site; expulsion of the rod;
 - 2. heavy vaginal bleeding that is unusual for this client;

- 3. concern that she might be pregnant, including delayed menstrual cycles after a long interval of regular cycles;
- onset or worsening of migraine headaches, repeated very painful headaches or blurred vision;
- 5. severe lower abdominal pain (rule out ectopic pregnancy).
- D. Management of Post-Insertion Side Effects/ Complications
 - 1. Arm pain, pus, or bleeding at insertion site
 - a. Management
 - (1) Advise the client to apply ice packs to the area for bruising, swelling, bleeding; moist heat for signs of infection.
 - (2) Advise to take Ibuprofen or other non-steroidal ant-inflammatory medication to relieve the discomfort.
 - (3) In case of infection of the insertion site, consultation with medical back-up may be indicated to select a therapeutic treatment drug.
 - b. Follow-up

Consider contacting the client within 48-72 hours to confirm improvement.

- c. Education
 - (1) Patient is instructed to keep wound site clean and dry for 24 hours.
 - (2) The patient should be informed that there might be irritation of a superficial nerve from the implants; paresthesia or paresthesia-like events may occur.
 - (3) Expulsion or migration of Implanon might be possible.
- 2. The implants appears to be coming out

Assessment/management - If the actual implant is protruding from the incision site, the implant should be removed and a new implant inserted at a different site.

- 3. Heavy or prolonged vaginal bleeding
 - a. Assessment
 - Review client history, including sexual history, other symptoms, contact to STD's.
 - (2) Physical examination and appropriate lab work should be done (according to Title X Guidelines) to rule out STDs.

- b. Management
 - (1) Any low-dose combination birth control pill for one or more cycles, if no contraindications to estrogen, or
 - (2) Ibuprofen 800 mg p.o. tid for 5-10 days, or
 - (3) Premarin 1.25 mg 2.5 mg p.o. qd for 20 days if no contraindications to estrogen.
- 4. Amenorrhea from the time of Implanon insertion, or after a pattern of regular periods
 - a. Assessment

Evaluate for pregnancy

- b. Management
 - (1) If pregnancy test is positive:
 - (a) Remove Implanon if client wishes to continue the pregnancy.
 - (b) Refer for immediate follow-up if ectopic pregnancy is suspected.
 - (c) Leave the Implanon in if the client plans an abortion.
 - (2) If the pregnancy test is negative:
 - (a) Discuss amenorrhea with client and reassure her that amenorrhea is a normal side effect of Implanon use.
 - (b) Implanon may be removed if client desires.
- 5. Severe lower abdominal pain
 - a. Assessment
 - (1) Client should be seen immediately to r/o pregnancy vs. PID vs. follicular cyst.
 - (2) Physical examination and appropriate lab work should be done (according to Title X Guidelines) to r/o pregnancy vs. PID vs. follicular cyst.
 - b. Management
 - (1) R/O pregnancy, PID, and/or follicular cyst.
 - (2) If unable to determine pathology, must be referred to physician for further evaluation.
- 6. Headache

- a. Assessment
 - (1) Review headache history.
 - (2) Take blood pressure.
- b. Management

Refer to physician for further evaluation, if indicated.

E. Client shall be advised to have an annual exam and Pap smear, based on the current Pap smear screening guidelines in use.

V. For Clients Desiring Removal

A. Subjective

If the client desires removal before three years, investigate the user's reasons for desiring removal. If, after counseling, the client still desires removal, the procedure should be scheduled.

B. Client Education

- The client needs to know that removal may take more time and may be more difficult than the insertion.
- 2. This information should be included in a removal consent.
- 3. The client should be counseled on all alternative contraceptive methods, and if she does not desire a pregnancy at this time, a method should be provided, as appropriate.

C. Follow-Up

Client should be encouraged to return for annual exam and Pap smear, based on the current Pap smear screening guidelines in use.

The following is a sample of an Implanon Insertion Consent Form. This form can be downloaded from the Women's Health Unit website at:

http://www.cdphe.state.co.us/pp/womens/FPNursingConsntsForms.html.

IMPLANON INSERTION CONSENT/ CLIENT INFORMATION SHEET

I understand that Implanon is a kind of birth control implant that consists of one small flexible rod, containing a form of the hormone progestin. It will be implanted just under the skin of my upper arm. I realize that Implanon will keep me from getting pregnant for 3 years. After three years, Implanon is no longer effective and must be taken out. I can have another Implanon inserted at that time or start using another method of birth control. I also know that I can have Implanon removed at any time and for any reason.

I understand that Implanon is more effective in preventing pregnancy than the Pill. Implanon is not permanent. I can get pregnant after it is taken out. I understand that Implanon might not be as effective if I take certain medications (mainly medications for seizures) or the herbal remedy called St. John's Wort.

I am aware that women who use Implanon have changes in their menstrual periods. Some women spot or bleed more often and some women bleed less or not at all. Spotting between periods is common. Periods can get more regular after 9-12 months. I understand that some women might also have a little weight gain, headaches or depression. I understand that Implanon does not protect me from sexually transmitted diseases, including the virus that causes AIDS (HIV). I need to use condoms, as well, to get protection from infections.

I have been told what to expect when Implanon is put in and taken out. I am aware that I might feel some discomfort during and after these procedures. I have been told about the problems that might occur when putting in or taking out Implanon such as: allergic response to anesthetic; bruising or soreness around the insertion site; or infection. After Implanon is put in, it could accidentally come out. When Implanon is taken out, it could break.

Removing Implanon may be slightly more difficult than putting it in. Sometimes it takes two clinic visits before the Implanon rod can be taken out.

I know to call the clinic if I:

Client Signature

- Want my Implanon taken out
- Have heavy bright red bleeding from the vagina that is more than a period
- Have a late period after my periods have been on time
- Have pain, pus, bleeding, or red skin where the Implanon was put in or if my Implanon capsule comes out
- Have very bad pain in the lower stomach or abdomen
- Have very bad headaches or problems with my vision

Based on my knowledge of the above, I consent to having Implanon inserted.

INTERPRETER' I have interpreted the information and advice present Implanon. I have also read to her the consent form ir content to her. To the best of my knowledge and beli consents to the insertion of Implanon.	ed orally to the client who has chosen to use a language she understands and explained its
Interpreter's signature	Date

Witness

Date

The following is a sample of Post Implanon Insertion Instructions. This form can be downloaded from the Women's Health Unit website at:

http://www.cdphe.state.co.us/pp/womens/FPNursingConsntsForms.html.

POST IMPLANON INSERTION INSTRUCTIONS

You can go back to normal daily activities immediately after the Implanon has been put in. After the numbness in your arm wears off, you may have some soreness for a day or two where the Implanon was inserted. There also may be some swelling, bruising, or discoloration for up to two weeks. This is how to care for your arm after Implanon is inserted. Please be aware of signs of infection and know how, when, and where to get medical care if needed.

- 1. Try not to bump the place where the Implanon was put in for a few days.
- 2. To make sure you don't get an infection where the Implanon was put in, keep the large gauze bandage on for 24 48 hours and keep it dry. Remove the large bandage after 24-48 hours.
- 3. Keep the little bandage strip on for 3 days, and keep it dry.
- 4. **If** you have any redness or oozing, or anything that concerns you, return to the clinic to have the insertion site checked.

After the incision has healed, you don't have to worry about bumping it or putting pressure on it. You can hold your child, carry books, do housework, or do whatever you usually do.

The Implanon is effective within 3 days if it was put in within 5 days of the first day of your period. You should use a back up method for _____ days.

The following is a sample of an Implanon Removal Consent Form. This form can be downloaded from the Women's Health Unit website at: http://www.cdphe.state.co.us/pp/womens/FPNursingConsntsForms.html.

Client Name		
IMPLANON REMOVAL CONSI	ENT/ CLIENT INFORMATION	
I am aware that if I don't want to get pregnant afte Implanon put in or choose a different method of bi		e a new
I understand that it could take up to 30 minutes to implant will be cleaned and numbed. Next, a sma so that it can be removed. I am aware that I migh	all cut will be made close to the tip	of the implant
I am aware of the possible problems that might of allergic reaction to the anesthetic; bruising or sore the implant could break; a second cut could be ne could be needed to take out the implant.	eness where the implant was remove	ved; infection;
Based on my knowledge of the above, I consent to	o the removal of Implanon.	
Client signature Date	Witness	Date
INTERPRETER'S	S STATEMENT	
I have interpreted the information and advictors to use Implanon. I have also read to her tand explained its content to her. To the best of mexplanation and voluntarily consents to the remove	he consent form in a language she y knowledge and belief she unders	e understands
Interpreter's signature		Date
POST IMPLANON REMOVAL INSTRUCTIONS I am aware that I can go back to my normal daily a removed. I might have some soreness for a day of also might be some swelling, bruising, or discolorate	or two where the Implanon was ren	
1. Try not to bump the place where the Implanor	was removed for a few days.	
2. Keep the large gauze bandage on for 24 - 48	hours and keep it dry to avoid an i	nfection

4. If there is any redness or oozing, or anything that concerns you, return to the clinic to have

3. Keep the little bandage strip on for 3 days, and keep it dry.

the removal site checked.

The following is a sample of a Hormonal Evaluation Form. This form can be downloaded from the Women's Health Unit website at:

http://www.cdphe.state.co.us/pp/womens/FPNursingConsntsForms.html.

HORMONAL METHOD EVALUATION ORAL CONTRACEPTIVES (Combined and POP), EVRA PATCH, NUVARING, IMPLANON (rod implant)	
Name	Today's date
Date of birth	Age
First day of last period	
1. Please check your current method:	
☐ Birth control pill (Combined)☐ Evra☐ Implanon	☐ Birth control pill (Progesterone only)☐ Nuvaring
Are you having any problems with you Explain:	_
3. Do you have any questions? ☐No Explain:	
4. Have you had any health problems or □No □?res Explain:	seen a physician since your last visit?
5. Are you taking any other medications List:6. Check if you have had any of the follow	
☐ Severe headaches	☐ Severe abdominal pain
□ Dizziness	□ Depression
Vision changes☐ Chest pain	☐ Nausea or vomiting☐ Heavy bleeding
Severe leg pain	☐ Weight gain
Client Signature	Date
TO BE COMPLETED BY STAFF	
S:	
O: B/P WT	
A :	
P:	